Three dimensional ultrasound of retinoblastoma: initial experience

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Aim: To use 3D ultrasonography (3DUS) for the diagnosis of retinoblastoma.

Methods: Five eyes of three children with retinoblastoma were evaluated using a commercially available computerised 3DUS system. Interactive sectioning of the stored and reconstructed 3D volumes were performed. 3DUS and histopathological findings were correlated after enucleation.

Results: 3DUS examination revealed characteristics consistent with retinoblastoma: endophytic mass, retinal detachment, intratumoral calcifications, and secondary orbital shadowing. Unlike 2D imaging, 3DUS allowed for analysis of the acquired and stored volumes. Rotation and sectioning of this volume allowed the discovery of new oblique and coronal views. For example, calcium related orbital shadows were seen as 3D volumes and (coronal) cross sections of the optic nerve were evaluated for evidence of intraneural invasion by retinoblastoma.

Conclusion: This is the first reported series of patients examined with 3DUS imaging for retinoblastoma. This technique allowed for new oblique and coronal views of the tumour and optic nerve. The ability to retrospectively analyse the stored ocular volume facilitated patient care, teaching, tumour-volume analysis, and telemedicine.

Retinoblastoma is the most common primary intraocular malignancy in children. Imaging techniques are essential for the diagnosis and follow up of retinoblastomas. Ultrasound has recently been used to reveal calcifications within the tumour and to evaluate extrascleral and optic nerve invasion. After treatment, ultrasonography and ophthalmoscopy are the main tools used to evaluate intraocular tumour regression.

Computer assisted three dimensional ultrasonography (3DUS) has become available to ophthalmological oncology. There is an emerging literature on the use of 3DUS for tumour diagnosis, radiation plaque placement, and follow up of treated choroidal melanomas.

This study presents the first reported use and our initial experience with 3DUS in evaluation of retinoblastoma. We have found new and exciting capabilities which have allowed us to produce unique and interesting views. Further experience and comparative studies will determine the role of 3DUS in the care of patients with retinoblastoma.

MATERIALS AND METHODS

Our methods of three dimensional ultrasonography (3DUS) have been described. In sum, we utilised a commercially available computerised 3D ultrasound system (Ophthalmic Technologies Incorporated (OTI) Downview, Ontario, Canada) to examine three patients (five eyes) with retinoblastoma. This system typically acquires data from 90 consecutive two dimensional (2D) images from which a 3D volume is rendered. The 3D volume is presented as a block, which was sliced, rotated, and examined from transverse, longitudinal, oblique, and coronal orientations.

RESULTS

Our first patient had a unilateral retinoblastoma. Before enucleation, 3DUS revealed multiple highly reflective nodules consistent with calcification (Fig 1A). Histopathology revealed a poorly differentiated retinoblastoma with intratumoral calcification and vitreous seeding (Fig 1B). Though focal extension to the optic nerve head was present ultrasound was not able to detect this small amount of optic nerve invasion. Tumour measurements from 3DUS were an apical height of 10.4 mm, a largest basal dimension of 14.9 mm, and a tumour volume of 909 mm³.

A 10 month old girl with bilateral retinoblastoma was examined with 3DUS which revealed a large intraocular tumour with multiple highly reflective calcifications. The optic nerve was sectioned for longitudinal, transverse, and unique coronal views. Where the tumour was found to overlay the optic nerve head, 3DUS allowed us to scroll through sequential coronal sections of the orbital portion of the optic nerve (Fig 1C). Interactive examination of the 3D volume revealed no evidence of intraneural tumour invasion or intraneural calcification. Histopathological examination revealed a poorly differentiated, retinoblastoma with focal necrosis, calcifications and no optic nerve invasion (Fig 1D).

Another 10 month old girl with bilateral retinoblastoma presented with a total retinal detachment, a large exophytic retinoblastoma, and a transretinal tumour (Fig 2A). With 3DUS, views clearly differentiated between the juxtapapillary tumour, adjacent retinal detachment, and the second trans-retinal tumour (Fig 2B). Not visible by ophthalmoscopy, the optic nerve was free of tumour by 3DUS (Fig 2B).

This patient’s right eye contained six non-macular tumours. Vitreous seeds could be imaged as overlying one of the tumours but could not be seen on ultrasound (Fig 2C). A unique coronal view allowed for three of the six tumours to be simultaneously imaged around the equator (Fig 2D). Intratumour calcification causes orbital shadowing which can be confused with the optic nerve. Interactive sectioning of the 3D volume allows the examiner to clearly demonstrate the optic nerve shadow (in one plane of section) while simultaneously imaging the orbital shadow in a second plane (Fig 2E and F). This technique (obtained by quartering the reconstructed ocular volume) cannot be done with 2D imaging.

DISCUSSION

Retinoblastomas are typically diagnosed before their third year of life. For the next 2 years, most ophthalmic
examinations are performed under anaesthesia. The risks of sedation requires the examiners be both quick and thorough.

We have found 3DUS to be uniquely valuable for evaluation of retinoblastomas. This is because 3DUS includes the ability to replay and reconstruct ultrasonograms in standard, oblique, and coronal views. Therefore, one main advantage of 3DUS (compared to 2DUS) is that it allows for multiple interactive reviews of reconstructed ocular volumes (without patient contact or anaesthesia). It also allowed for the first ultrasound generated coronal views of the optic nerve.

These unique 3DUS capabilities become even more important when one considers the findings that have been associated with an increased risk for metastasis: tumour contact with the choroid, optic nerve invasion, and extrascleral extension. Unlike standard two dimensional ultrasound (2DUS), 3DUS offers the potential to quantify the area of tumour contact with the choroid. With 3DUS, extrascleral extension and optic nerve invasion can be scrutinised with unique previously unavailable oblique and coronal sections. Though none of our patients exhibited intraneural tumour invasion, 3DUS clearly allowed for the first ultrasonographic views of serially examined coronal sections of the optic nerve.

Other unique findings included calcium related 3D orbital shadowing, unique coronal sections of the calcified tumour, as well as simultaneous views of a cross section the optic nerve and the tumour (obtained by quartering the reconstructed ocular volume). These cases demonstrate that 3DUS offers unique and valuable capabilities that can be used to improve the management of retinoblastoma.

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REFERENCES